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(54) Title: METHOD FOR DETECTING POINT MUTATIONS IN DNA UTILIZING FLUORESCENCE ENERGY TRANSFER

(57) Abstract

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A method for detecting point mutations in DNA using a fluorescently labeled oligomeric probe and fluorescence resonance energy transfer (FRET) is disclosed. The selected probe is initially labeled at each end with a fluorescence dye, which act together as a donor/acceptor pair for FRET. The fluorescence emission from the dyes changes dramatically from the duplex stage, wherein the probe is hybridized to the complementary strand of DNA, to the single strand stage, when the probe is melted to become detached from the DNA. The change in fluorescence is caused by the dyes coming into closer proximity after melting occurs and the probe becomes detached from the DNA strand. The change in fluorescence emission as a function of temperature is used to calculate the melting temperature of the complex or T_m. In the case where there is a base mismatch between the probe and the DNA strand, indicating a point mutation, the T_m has been found to be significantly lower than the T_m for a perfectly match probe/target duplex. The present invention allows for the detection of the existence and magnitude of T_m, which allows for the quick and accurate detection of a point mutation in the DNA strand and, in some applications, the determination of the approximate location of the mutation within the sequence.

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METHOD FOR DETECTING POINT MUTATIONS IN DNA UTILIZING FLUORESCENCE ENERGY TRANSFER

This invention relates to the identification of genetic mutations and, more particularly, to a method of detecting point mutations in DNA utilizing fluorescence energy transfer. This invention was supported by National Institutes for Health grant DK36288 to the Center for Biotechnology, University of Nebraska-Lincoln.

Background

The detection of mutations in sequences of DNA is becoming increasingly important in medical science. The detection of such a mutation in a DNA sequence typically involves the use of an oligodeoxyribonucleotide probe that is complementary to the target DNA sequence. The probe is designed to present some moiety, such as a radioactive element, that signals the occurrence of hybridization in a filter assay or an electrophoretic gel. The identification of hybridization has been used diagnostically for specific bacterial infections by detection of *Mycobacterium tuberculosis* genomic DNA, gonorrhea rRNA, *Chlamydia* genomic and plasmid DNA and *Escherichia coli* and *Bacillus subtilis* rRNA. Hybridization assays have also been developed for viral detection, including cytomegalovirus (CMV), human papilloma virus (HPV), and HIV-1.

By combining target amplification with allele specific oligonucleotides, small samples of human DNA can be analyzed for purposes of genetic screening, including the study of genetic changes associated with well-known inherited diseases. For instance, cancers typically display familial site-specific clustering. The identification of this kind of clustering can aid in the determination of enhanced risk for the development of the particular cancer. In addition, hereditary metabolic variations in DNA have been identified that affect the metabolism of known carcinogens. A variation that would increase the

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metabolism of a carcinogen may impact the likelihood of the development of cancer and, if developed, the speed of the cancer's growth.

Traditional hybridization methods have been developed which employ radioactive probes with separation on filters. While radioactive probes have performed suitably well, growing concern over the use of radioactive materials has stimulated a search for alternative probes that achieve similar levels of sensitivity and performance without the risks and dangers associated with radioactive materials. For instance, biotin has been incorporated into an oligodeoxyribonucleotide for use in biotinavidin-linked analyses. In addition, numerous modifications of DNA have been used in the development of other alternative probes, including links to antibodies, gold-antibodies, mercury for double antibody reactions, eupsoralen, and fluorescent dye links for fluorescence detection of hybridization. These alternative methods typically allow approximately 10⁵ to 10⁶ copies of the DNA to be detected.

These and related advancements in the art have given rise to several methods of DNA mutation detection. These methods include denaturing gradient gel electrophoresis (DGGE), single-strand conformational polymorphisms (SSCP), temperature gradient gel electrophoresis (TGGE), the heteroduplex method (HET), ribonuclease cleavage, chemical cleavage of mismatch (CCU), ligase assay, allelespecific amplification (ASA) dideoxy fingerprinting (ddF), and allele-specific oligonucleotides (ASO). DGGE, SSCP, TGGE, HET, and ddF are frequently used to locate which exons of a gene contain mutations.

The currently available non-radioactive methods for detecting mutations in DNA have been somewhat problematic. For example, these methods have been generally unable to consistently provide accurate results in detecting point mutations in DNA. These detection methods have also proven to be time-consuming and quite costly to use. In addition, these non-radioactive mechanisms require a significant amount of DNA to perform their detecting function, though many times only a small quantity of DNA is available for analysis. Moreover,

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these methods are difficult to use, often requiring complex instruments and highly trained technicians not available in many laboratories. Finally, the materials utilized in these methods are generally either fragile or prone to degradation during the testing procedure.

Summary of the Invention

It is, therefore, a primary object of the present invention to provide a method for quickly and accurately detecting point mutations in DNA.

It is also an object of the present invention to provide a method for quickly and accurately detecting point mutations in DNA that does not utilize radioactive material.

It is a further object of the present invention to provide a method for quickly and accurately detecting point mutations in DNA that requires only a relatively small amount of DNA to perform its detection function.

It is another object of the present invention to provide a method for quickly and accurately detecting point mutations in DNA that is inexpensive, noncomplex and easy to use.

It is yet another object of the present invention to provide a method for quickly and detecting point mutations in DNA that is durable and less likely to experience degradation of its constituent components during the testing procedure.

It is yet another object of the present invention to provide a method for deteting point mutations in DNA which requires minimal sample preparation.

To accomplish these and related objectives, the present invention relates to a method for detecting point mutations in DNA using a fluorescently labeled oligomeric probe and fluorescence resonance energy transfer (FRET). The selected probe is labeled at each end with a fluorescent dye, which act together as a donor/acceptor pair for FRET. The fluorescence emission from the dyes changes dramatically from a probe/target duplex stage, wherein the probe is

hybridized to the complementary strand of target DNA, to the single strand stage, when the probe is melted to become detached from the target DNA. The change in fluorescence is caused by the dyes coming into closer proximity after melting occurs and the probe becomes detached from the target DNA strand. The change in fluorescence emission as a function of temperature is used to calculate the melting temperature of the complex or T_m . Where there is a base mismatch between the probe and the target DNA strand, indicating a point mutation in the target DNA strand, the T_m has been found to be significantly lower than the T_m for a perfectly match probe/target duplex. The present invention allows for the detection of the T_m , which allows for the quick and accurate detection of a point mutation in the target DNA strand.

Brief Description of the Drawings

Fig. 1 illustrates the structure of a representative 16-base oligomer probe for use in connection with the method of the present invention. The probe includes x-rhodamine attached to the 5' end of the probe and fluorescein attached to the 3' end, which act as a acceptor and donor, respectively, for FRET;

Fig. 2 is a graph plotting fluorescence intensity versus wavelength and showing steady-state fluorescence emission scans for the probe of Fig. 1 as a hybridized duplex structure with its complementary target DNA strand in the solid curve, and as a single strand upon melting in the broken curve;

Fig. 3 is a graph plotting the fraction of probe hybridized to target DNA versus temperature and showing melting curves for a perfectly matched duplex of the probe of Fig. 1 and its complementary target DNA strand, and the probe of Fig. 1 with its complementary target DNA strand having a single central base substitution, the vertical broken lines indicating the melting point for each respective curve; and

Fig. 4 is a is a graph plotting the fraction of probe hybridized to target DNA versus temperature and showing the melting curve for a duplex

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including the probe of Fig. 1 bound to a 7249-base phage target DNA, M13mp18+, which contains a 16-base complementary sequence, the vertical broken line indicating the melting point of the duplex.

Description of the Preferred Embodiment

The method of the present invention is carried out utilizing a probe, which is typically a deoxyribonucleotide oligomer. An illustrative 16-base probe is shown in Fig. 1. The base length and specific nucleotide sequence of the probe will, of course, vary depending on the target sequence of the DNA strand. A probe having a length of 10 to 30 bases have been found to work suitably well. The probe length must be sufficient for essentially full hybridization so an accurate melting transition can be detected even for the mismatch. It is understood that the precise sequence of the probe may be altered or engineered to complementarily match the target DNA strand to be assayed and, consequently, all such sequences are within the scope of the present invention. The probe can be used in a solution or on a solid-phase support, depending on the application.

The probe of the present invention must be fluorescently labeled. A pair of dyes are covalently linked to the probe as labeling agents. The dyes act as a donor/acceptor pair for FRET. The efficiency of the energy transfer from the donor dye to the acceptor dye is sensitive to distance, the efficiency rising dramatically as the distance between the probes is decreased. When the probe is attached to the target DNA sequence, the dyes are relatively spaced apart, thereby inhibiting efficient transfer of energy between the dyes. When the probe becomes detached from the DNA, the dyes come into closer proximity. The energy transfer between the dyes rises, which is indicative of the melting of the duplex.

A number of considerations are involved in choosing a donor/acceptor dye pair and in designing the probe. The dyes should have a high molar absorptivity and quantum yield, and the fluorescence lifetime of the donor should be sufficiently long to ensure accurate measurement. The emission of the

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dyes should have limited sensitivity to factors other than changes in distance. For instance, the probe should be resistant to changes in pH, ion concentration and to photobleaching. Another important consideration is the value of R_0 (the distance at which FRET is 50% efficient), relative to the length of the oligomer. The value of R₀ must therefore be calculated or at least estimated. This value derives from the overlap of the emission band of the donor dye with the absorption band of the acceptor dye, and also from the region of the wavelength range on which the overlap integral is centered. Further, the dyes must be attached to the oligomer with sufficient freedom of rotation so that the average value of the orientation factor K2, will be close to two-thirds. With these considerations taken into account, numerous successful probes have been designed ranging from 14 - 20 bases and with the following donor/acceptor pairs: fluorescein/tetramethylrhodamine, fluorescein/x-rhodamine, and fluorescein/Cy-3. In all of these probes, the linker arms between the oligomer and the dyes were either 3- or 6-carbon chains. It is to be understood, however, that other probes and donor/acceptor pairs may be developed in accordance with this invention and are, thus, within its scope.

The difference in the fluorescence emission of the probe as a single strand or in a duplex structure derives from a difference in an excitonic interaction in the weak-coupling limit, an interaction that varies with the inverse sixth power of the distance between transition moment dipoles of the donor and acceptor dyes or moeties. The steady-state fluorescence emission intensity of either dye is proportional to its quantum yield, which, neglecting static quenching, is the ratio of the rate constant for fluorescence (k_F) , which derives from the Einstein B coefficient, to the sum of the microscopic rate constants that deactivate the state from which the fluorescence arises, which is assumed to be the lowest singlet (S_1) . These processes can be summed as follows: $k_F + k_I + k_I$; where k_I is the sum of all nonfluorescence processes that deactivate S_1 in the absence of acceptor, and k_I is

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the rate constant for resonance energy transfer. The quantum yield of donor fluorescence in the presence of acceptor is then:

$$\Phi_{t} = k_{F}/(k_{F} + k_{I} + k_{t}) = k_{F}/[(k_{F} + k_{I})(1 + X^{6})]$$

$$= \Phi_{O}/(1 + X^{6}),$$

where $\Phi_{\rm O}$ is the quantum yield in the absence of transfer or, in other words, the absence of acceptor. The quantity X is $R_{\rm O}/R$, where R is the distance between the donor and acceptor, and $R_{\rm O}$ is the donor/acceptor distance at which the rate constant for transfer equals $k_{\rm F} + k_{\rm I}$. $R_{\rm O}$ can be calculated relatively easily by one skilled in the art from the measured spectroscopic properties of the donor and acceptor and applying the assumptions with respect to the effective refractive index of the medium separating the donor and acceptor and the rotational averaging of the angular portion of the dipole-dipole interaction. It is assumed that $R_{\rm O}$ for fluorescing/x-rhodamine (60 Å) remains constant throughout the melting process and, therefore, changes in the steady-state intensity, which are proportional to $\Phi_{\rm D}$ derive from changes in R.

The donor and acceptor dyes may be attached to the probe at any two points. It is preferred that the dyes be spaced relatively far apart on the probe. Most preferably, the dyes are attached at opposite ends of the probe. Because the FRET donor and acceptor are preferably relatively far apart, FRET is minimized (k_t is low) for a probe/target DNA duplex. When the dyes are spaced apart, the donor emission is high and acceptor emission is low. This is illustrated by the solid line in Fig. 2. Upon melting to two single strands, the duplex structure is lost and more flexible single strand brings the donor and acceptor closer together. Consequently, FRET is significantly increased, and donor emission decreases and acceptor emission increases. This is shown by the broken line in Fig. 2. The melting temperature, T_m , is the temperature at which the fraction (f) of the total probe bound to complement is 0.5. In another embodiment, the FRET acceptor acts as a quencher having its own fluorescence emission.

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The melting process is a sufficiently sharp transition that it may be treated as a two-state process. This can be expressed mathematically by considering B to be the equilibrium concentration of B DNA, and C_T and P_T , respectively, to be the total target and probe concentrations. The equilibrium constant for helix formation is then $K = B / [C_T - B) (C_P - B)]$ and $f = B / P_T$, with all concentrations referenced to a standard state of 1 M. The equilibrium constant for $C_T \ge P_T$ can then be written as follows:

$$K = [f/(1-f)][P_T(1-f) + (C_T - P_T)]^{-1}.$$
with $[f/(1-f)] = 1$ at T_m .

10 At T_m , $\triangle G = 0$, and

$$\Delta G^{\circ} = -(\mathbf{R}T_m)\ln(K),$$

where **R** is the molar gas constant. If values of $\triangle H^{\circ}$ and $\triangle S^{\circ}$ are available for the given solvent conditions and assumed to be independent of temperature up to T_m then, since $\triangle G^{\circ} = \triangle H^{\circ} - T_m \triangle S^{\circ}$, T_m can be calculated from this formula:

$$T_m = \Delta H^{\circ}/[\Delta S - \mathbf{R} \ln(K)],$$

This calculation is dependent upon probe and target concentrations through K, and the steepness of the transition at the midpoint, $\partial f/\partial T$, gives ΔH° at T_m for a true two-state process, and it can be shown that

$$\Delta H^{\circ} = \mathbf{R} T_{\mathrm{m}}^{2} (\partial f / \partial T) [4 + 1/(C_{T} / P_{T} - 1/2)].$$

20 Alternatively, T_m can be predicted from empirical equations.

The ΔT_m for a DNA sequence due to a point mutation or base-pair mismatch has been found to be from about 2 degrees to about 12.5°C as compared to a perfect probe/target match. The ΔT_m has been found to be dependent on the length of the probe/DNA strand duplex and also upon the location of the point mutation or base-pair mismatch in the duplex. For instance, if the point mutation lies toward either end of the duplex, ΔT_m will generally be lower, typically in the range of about 6 to about 8°C. If the point mutation or base-pair mismatch resides toward the center of the duplex, ΔT_m will be relatively higher, typically in the range of about 9 to about 11°C, it has been found that ΔT_m is generally equivalent

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to about 1 to about 1.5°C per percent base mismatch in the oligomer probe. It is to be appreciated that ΔT_m will vary depending on not only the length of the duplex and location of mutation, but also will vary depending on the specific order of the sequence. Consequently, all such variances are within the scope of the disclosure.

EXAMPLE

The following example illustrates the performance of the method of the present invention. The example should be construed only as illustrative and not in limiting sense.

For purposes of this example, a 16-base deoxyribonucleotide oligomer with x-rhodamine covalently bound to the 5' end and fluorescein to the 3' end will be discussed. This probe is shown in Fig. 1 and will be addressed as R*16*F. As seen in Fig. 1, the R*16*F probe comprises x-rhodamine-5'-GTAAAACGACGGCCAG-3'-fluorescein. The R*16*F probe was designed to be complementary to bases 6291 to 6306 of the 7249-base phage DNA M13mp18(+), one base removed from the restriction site. Hybridization of the R*16*F to a complementary strand results in a B DNA helical structure with the 3'-fluorescein and the 5'-x-rhodamine positioned at opposite ends.

The R*16*F probe was selected with several considerations in mind. First, the length of the probe is sufficient for the melting temperature of the duplex to be above room temperature, ensuring complete hybridization at 25°C. Second, for Förster energy transfer to reflect small changes in interdye distances, the average distance apart should be near R_{θ} . The average distance in the duplex would be approximately 10% greater than R_{θ} . Finally, the placement of dyes on bases 1 and 16 results in their being located 180 degrees apart when viewed down the helix axis, enhancing their separation and the signal change associated with melting.

A steady-state fluorimeter with computerized data acquisition (Photon Technology International, Inc., Model A-1010) was modified to use as

excitation light the 488-nm line of a Coherent Innova 70-4 Ar laser. A controllable shutter between the excitation beam and the cuvette was utilized to ensure that the solution was exposed to excitation light only during the course of a scan, to minimize the photobleaching of fluorescein. The R*16*F probe was obtained from Research Genetics in Huntsville, Alabama. The perfect (CTGGCCGTCGTTTTAC) and single mismatch (CTGGCCGTTGTTTAC) complements were synthesized by the inventors, and the M13mp18(+) was from obtained Sigma in St. Louis, Missouri.

The buffer was $0.01 \, M$ NaPi, pH 8, 1 mM EDTA and $0.18 \, M$ NaCI, with a total volume of $250 \mu l$ in a stoppered microcuvette. Stock solutions of R*16*F and its complement were diluted to 10 and 20 nM, respectively, and allowed to hybridize for 45 minutes, essentially to completion. The duplex with the mismatch was monitored spectroscopically for the last 15 minutes of the procedure to ensure that no further change occurred.

For the R*16*F/M13 duplex, stock solutions were diluted to 6.6 and 13.2 nM, respectively, and the solution left for 2.5 minutes in a heat block (Thermodyne, model 17600) at 65°C. Hybridization occurred as the solution returned slowly to 20°C over 1 hour. The cuvette was capped prior to data collection to prevent evaporation at high temperatures. Scans were collected from 502 to 625 nm every 3 to 4°C, from 16 to 74.5°C, 16 to 67°C, and 20 to 76°C for the R*16*F duplex, mismatch duplex, and M13 duplex, respectively. The solutions were equilibrated for 10 minutes at each temperature prior to scanning.

The temperature of the solution was controlled using flow through the cuvette holder from a Lauda K-2/R constant-temperature circulating bath. The temperature of the circulating water for each scan was noted. After the experiment, a thermistor was placed in the buffer-filled cuvette to obtain the actual solution temperature for each temperature noted previously. Baseline scans, which included the 583nm Raman scattering from water and background introduced with

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the M13, were collected prior to each data set and subtracted from all subsequent scans prior to analysis.

The emission scan of the R*16*F contained two peaks: The peak at 520 nm was derived from direct absorption by fluorescein of the 488-nm excitation light, modified by FRET. The peak at 610 nm was due primarily to FRET from the fluorescein to the x-rhodamine, with a small contribution from direct absorbance by x-rhodamine of the excitation light. In the perferred embodiment, the fraction, f, of R*16*F proble hybridized to its complement target DNA is represented in the following equation:

$$f = \delta(Y) / [1 - Y(1 - \delta)],$$

where

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$$\delta = I_{\text{rhod}}^{()}/I_{\text{rhod}}^{H}$$

and

$$Y = (\langle p \rangle - p^{U}) / (p^{H} - p^{U}),$$

and I_{rhod} is the peak rhodamine emission intensity, p is the ratio of the 520 to 610 nm peaks, $\langle p \rangle$ is any observed intermediate value of p, and the superscripts U and H refer, respectively, to unhybridized and hybridized R*16*F. The fraction of bound R*16*F was calculated at each temperature measured, and was used to construct the melting curves. The sets of data points on each plot that correspond to the premelting and postmelting regions and to the melting process were each fit by linear regression and the T_m taken as the mid-point of the transition region.

In another embodiment, a fraction of the probe hybridized to target DNA is determined by analyzing the donor emission intensity in unhybridized and hybridized states. This embodiment enables a non-fluorescent quencher to be used as the accepted dye, although it does not require that the acceptor be non-fluorescent.

The fluorescence emission scans of R*16*F differ dramatically for the duplexed and single-strand forms, as discussed above in Fig. 2. Melting results in a decrease in the fluorescence emission of the fluorescein and an increase in the

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rhodamine emission, which is the result of the decrease in the inter-dye distance. Fig. 3 illustrates melting curves for the duplexes formed by R*16*F hybridized to the perfect 16-base complement and to the 16-base complement with a single C \rightarrow T base substitution. The T_m s for these duplexes are 58.3 and 48.2°C, respectively. Fig. 4 shows the melting curve for the duplex formed between R*16*F and its target sequence in M13mp18(+), with a T_m of 56.4°C. The mismatch is easily detected and gives rise to a ΔT_m of 10°C.

The T_m of the perfect complement can be predicted rather well from this empirical relationship:

T_m(°C)= $81.5+16.6(log_{10}[Na+])+0.41(%G+u.c.)-600/N$,

where N is the number of bases in the shorter oligo strand. For the R*16*F probe, the predicted T_m of 55.5°C is consistent with the observed values of 56.4 and 58.3°C for M13 and the 16-mer complement, respectively.

 T_m is expected to be insensitive to secondary or higher structures around the probe site which are known by kinetic studies to characterize M13mp18(+). A comparison of Figs. 3 and 4 shows that T_m is, in fact, nearly insensitive to the length of the target DNA, particularly considering that the target and probe concentrations are lower for the M13. Assuming that the standard state enthalpy and entropy changes are the same for the two duplexes, a lower T_m may be predicted for the M13 by about 0.9 degrees. It is not known whether the remaining 1-degree difference is insignificant or reflects a perturbation of the target structure. More involved calculations of the melting temperature can be made in which nearest-neighbor interactions are included in ΔH° and ΔS° . However, these quantities are not available for the above buffer conditions, and the above empirical relationship provides a sufficiently close estimate of T_m for the purpose of experimental design.

The following thermodynamic quantities for the probe hybridized to the 16-base complement were calculated: $\Delta G^{\circ} = -49.6$ kJ/mole, $\Delta H^{\circ} = -373$ ± 21 kJ/mole of duplex (-23.3 kJ/mole of base pair) and $\Delta S^{\circ} = -977 \pm 17$ J/mole-K

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at $T_m = 331.5$ K. Using values from Breslauer et al. for the disruption of a CG base pair adjacent to a CG base pair (ΔH° and ΔS° more positive by about 50 kJ/mole and 116 J/mole-K), a T_m of 320.1 K for the mismatched duplex can be calculated. This is a decrease in T_m of 11.4°C, which is in agreement with the 10.1°C decrease observed. In this calculation, the changes in ΔH° and ΔS° for lowering [Na+] from 1 M to 0.2 M have been neglected.

It is convenient to have measurable fluorescence for both the doublelabeled oligo as a single strand and in the duplex, as well as sensitivity of the fluorescence to changes in the donor/acceptor distance. If θ as the ratio of the quantum yield in the presence of the acceptor to that in the absence, then $\theta = (1 - 1)^{-1}$ $+X^{6}$)⁻¹ = 1 - η , where η is the efficiency of transfer. It is straightforward to show that the maximum sensitivity of θ , where sensitivity is defined as $-d\theta/dX$, to a change in R occurs at $R = R^* = (0.946)R_0$, for which $-d\theta/dX - 1.54$. For $R = R_0$, $-d\theta/dX = 1.5$, and for either R* or R₀, a 1% change in fluorescence intensity corresponds to a change of 0.6Å in R for $R_0 = 60$ Å. The sensitivity decreases in 1 on either side of R_0 (at R = 1.344 R_0 and at R = 0.847 R_0) where the respective values of θ are 0.855 and 0.269, a normalized change from 1 to 0.32. For the 16-base oligomer, distance changes on melting correspond to $R = 1.142 R_0$, decreasing to $R = 0.858 R_0$, and the change in normalized fluorescence intensity for the donor, as in Fig. 2, is from 1 to 0.44. These distance considerations are linked to the desired ΔT_m , which increases as the length of the probe decreases and is maximized for location of the mismatch at or near the center of the probe.

Both optical and calorimetric measurements of the melting transition agree for a 0-mer duplex havingGC base pairs on the ends, showing that the process can be treated to high precision as a two-state process. Intermediates have been detected for single strand melting, however, and there may be instances where bases, particularly at the ends of the duplex, are unpaired before full strand separation. On the other hand, the large change in fluorescence emission reported here would be expected only after actual strand separation and, thus, might be

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expected to lead to the appearance of a sharper transition than that observed by absorbance. Monitoring absorbance changes associated with melting is not feasible for duplexes between short oligomers and long strands of DNA due to the overwhelming background from the long strand.

The long DNA strand tested in this example was single-stranded. Special hybridization techniques are required with DNA that is originally duplex. The melting process, once the probe has bound, should closely resemble that of the example. It is essentially independent of higher order structure in the vicinity of the target, owing to the large ΔH° of melting. In practice, one would titrate the probe with DNA to determine the number of copies of the target, and then carry out the melting experiments as described above. Once hybridization is achieved and having an estimate of T_m from the foregoing equations, one could start the temperature ramp about 10 degrees below the expected T_m for the mismatch and complete the analysis within 1 hour.

The sharp transition detected by FRET using the double-labeled oligonucleotide allows a rapid and clear determination of whether a mismatch is present in the target. The quickness and accuracy of this method are not found in methods and procedures currently available in the art. Further, the method of the present invention is straightforward with few steps to follow. A technician will be able to carry out this method quite easily with little training and, thus, the method is economical and efficient. The materials used in this method are hearty and unlikely to degrade or break during the procedure. This will reduce the need to repeatedly perform the testing due to failure of the components. In addition, only a small amount of DNA is needed to make a point mutation determination utilizing this method. The ultimate beneficiary of the method of the present invention will be the patient, who will obtain results quickly and inexpensively and who will be able to rely upon the accuracy and reliability of the results of the test.

It is, therefore, readily apparent that this invention is capable of attaining the objects set forth above as well as other advantages that are obvious

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and inherent to the invention. It is to be understood that other subcombinations are of utility and may be employed without reference to other features and combinations of the invention. Because many embodiments may be made of the invention without departing from the scope of this disclosure, it is to be understood that all matter set forth herein is to be interpreted as illustrative only.

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WHAT IS CLAIMED IS:

- DNA sequence, wherein the sequence has a reference melting point at which it becomes detached from a perfectly matched complementary sequence, the method comprising: providing a probe having a base sequence complementary to the target DNA sequence; attaching two dyes to the probe, the dyes acting together as a donor/acceptor pair for FRET and producing FRET and producing FRET emissions; allowing the probe to hybridize with the DNA sequence to form a probe/target DNA duplex; recording an attached value of the FRET emissions from the dyes of the probe/DNA duplex; heating the probe/DNA duplex; monitoring the temperature at which the FRET emissions from the probe and duplex are reduced to a detached value; and determining an actual melting point of the probe/target DNA duplex by recording the temperature of the probe/DNA duplex at the midpoint between the attached value of the FRET emissions and the detached value of the FRET emissions; whereby if the actual melting point is lower than the reference melting point a point mutation exists.
- 2. The method of claim 1 wherein the dyes are attached to opposite ends of the probe.
- 3. The method of claim 2 whereby a point mutation exists if the actual melting point is about 2° to about 12.5°C lower than the reference melting point.
 - 4. The method of claim 3 whereby a point mutation exists if the actual melting point is about 6 to about 11°C lower than the reference melting point.
- 5. The method of claim 4 wherein the probe is an oligodeoxyribonucleotide.
 - 6. The method of claim 5 wherein the probe is a oligomer having a length of 10 to 30 bases.

- DNA sequence, wherein the sequence has a reference melting point at which it becomes detached from a perfectly matched complementary sequence, the method comprising: supplying a probe having a base sequence complementary to the target DNA sequence and being labeled with at least one dye; allowing the probe and the DNA sequence to hybridize to form a probe/DNA duplex; heating the probe/DNA duplex; determining an actual melting point for the probe/DNA duplex by detecting a change in fluorescence emitted by the dye; assessing whether a point mutation exists in the target DNA sequence by comparing the reference melting point of the sequence to the actual melting point of the duplex; whereby if the actual melting point is lower than the reference melting point, a point mutation exists.
- 8. The method of claim 7 wherein at least two dyes are attached to the probe.
- 9. The method of claim 8 wherein the dyes are attached to opposite ends of the probe.
 - 10. The method of claim 9 whereby a mutation exists if the actual melting point is from about 2° to about 12.5°C lower than the reference melting point.
- 11. The method of claim 10 whereby a point mutation exists if 20 the actual melting point is from about 6° to about 11°C lower than the reference melting point.

- 12. A method for detecting the location of a point mutation in a selected target DNA sequence, wherein the sequence has a reference melting point at which it becomes detached from a perfectly matched complementary sequence, the method comprising: providing a probe having a base sequence complementary to the target DNA sequence; attaching two fluorescent dyes to the probe, the dyes acting together as a donor/acceptor pair for FRET and producing FRET emissions: allowing the probe to hybridize with the DNA sequence to form a probe/DNA duplex; recording an attached value of the FRET emissions from the dyes of the probe/DNA duplex; heating the probe/DNA duplex; monitoring the temperature at which the FRET emissions from the probe/DNA duplex are reduced to a detached value; and determining an actual melting point of the probe/DNA duplex by recording the temperature of the probe/DNA duplex at the midpoint between the attached value of the FRET emissions and the detached value of the FRET emissions; whereby if the actual melting point is about 9 to about 11°C lower than the reference melting point, the point mutation is located near the center of the target DNA sequence, and whereby if the actual melting point is about 6 to about 9°C lower than the reference melting point, the point mutation is not located near the center of the target DNA sequence.
- 13. The method of claim 12 wherein the dyes are attached to opposite ends of the probe.
 - 14. The method of claim 13 wherein the probe is an oligodeoxyribonucleotide.
 - 15. The method of claim 14 wherein the probe is a oligomer having a length of 10 to 30 bases.
- 25 If. The method of claim 15 wherein the probe is a 16-base oligomer.

X-RHODAMINE

FIG. 1

FLUORESCEIN

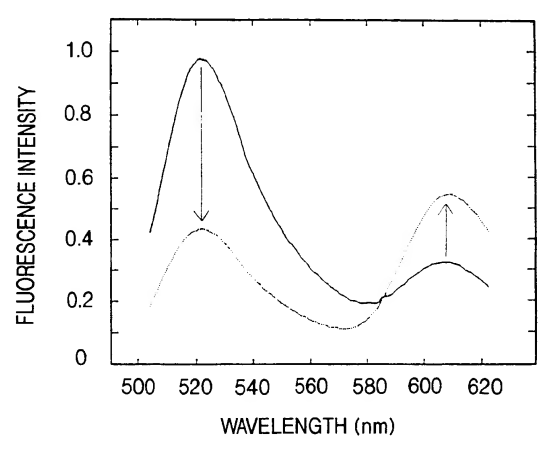


FIG. 2

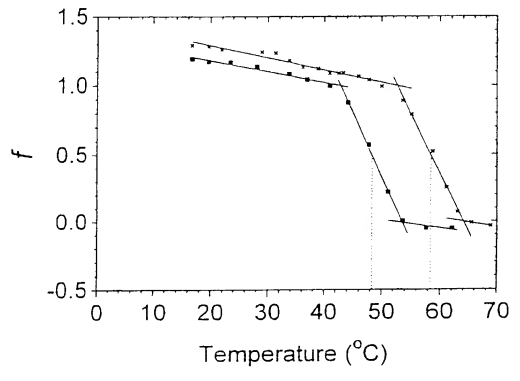
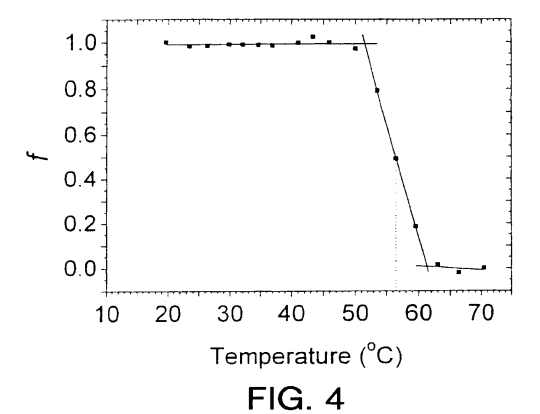


FIG. 3



SUBSTITUTE SHEET (RULE 26)

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/19525

| IPC(6) | SSIFICATION OF SUBJECT MATTER :C12Q 1/68 | | | |
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| US CL According | : 435/6 to International Patent Classification (IPC) or to both | national classification and IPC | | |
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| Minimum d | locumentation searched (classification system follow | ed by classification symbols) | | |
| U.S. : | 435/6; 536/24.3 | | | |
| Documenta | tion searched other than minimum documentation to the | he extent that such documents are included | in the fields searched | |
| i | data base consulted during the international search (ree Extra Sheet. | name of data base and, where practicable | , search terms used) | |
| C. DOC | CUMENTS CONSIDERED TO BE RELEVANT | | | |
| Category* | Citation of document, with indication, where a | ppropriate, of the relevant passages | Relevant to claim No. | |
| A | CARDULLO et al. Detection of r nonradiative fluorescence resonance e the National Academy of Sciences, U. No. 23, pages 8790-8794, especially | nergy transfer. Proceedings of SA. December 1998, Vol. 85, | 1-16 | |
| Α | UCHIYAMA et al. Detection of Ur Vivo by Fluorescence Resonance End Biological Chemistry. January 1996, 384, especially abstract. | ergy Transfer. The Journal of | 1-16 | |
| X Further documents are listed in the continuation of Box C. See patent family annex. | | | | |
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| Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Authorized office DEBRA SHOEMAKER | | | WW 19 | |
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| Category* | Citation of document, with indication, where appropriate, of the relevant passages | Relevant to claim No | | |
| Y | IITIA et al. Detection of a Point Mutation Using Short Oligonucleotide Probes in Allele-Specific Hybridization. BioTechniques. September 1994, Vol. 17, No. 3 pages 566-573, especially Figure 1. | 1-16 | | |
| Y, P | US 5,591,578 A (MEADE et al.) 07 June 1997, especially Figure 1. | 1-16 | | |
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/19525

| B. FIELDS SEARCHED Electronic data bases consulted (Name of data base and where practicable terms used): | | | | | |
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| Searched inventors and keywords: fluroescence energy transfer or fret and donor or and probe or oligonucleotide and detect with mismatch or mutation in APS, CAPLUS EMBASE, BIOSIS and WPIDS. | Searched inventors and keywords: fluroescence energy transfer or fret and donor or quencher or two and same or one and probe or oligonucleotide and detect with mismatch or mutation in APS, CAPLUS, MEDLINE, SCISEARCH, EMBASE, BIOSIS and WPIDS. | | | | |
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